Synthesis, Characterization and Antibacterial Activity of Some Amino Acid Derivatives

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ABSTRACT

The present work includes the synthesis of glycine and L-amino acid derivatives 6-10 (A,B) via Schiff’s bases 3 (A,B), which were obtained from the reaction of 2-aminopyridine 2 with benzaldehyde 1A or 4-chlorobenzaldehyde 1B. The reaction of 3 (A,B) with benzoyl chloride yielded benzamide derivatives 5 (A,B). The synthesis of 6-10 (A,B) has been performed by the reaction of 5 (A,B) with (glycine, L-alanine, L-phenylalanine, L-aspartic acid and L-asparagine). Infrared and nuclear magnetic spectroscopic techniques Fourier Transform Infrared (FT-IR), Proton Nuclear Magnetic Resonance (¹H-NMR) and Carbon-13 nuclear magnetic resonance (¹³C-NMR) were used to characterize the newly synthesized compounds. The antibacterial activity of final products has been evaluated against two kinds of Gram-positive and Gram-negative bacteria (Staphylococcus aureus and Klebsiella pneumonia). Overall, results indicate a lower and moderate antibacterial activity in comparison with meropenem as a reference against both bacteria. The highest activity was obtained using the compound 7B for both kinds of bacteria and 6A for only K. pneumonia.

Keywords: Schiff’s bases, 2-aminopyridine, L-amino acids, Antibacterial activity, Benzamide

INTRODUCTION

In the last decade, Schiff’s bases have been indicated as anti-proliferative [1], anti-inflammatory [2], antimicrobial [3], antiviral [4], antimalarial [5], anticancer [6], antibacterial [7], antifungal [8] and antitumor [9] active compounds. Schiff’s bases can be considered as wide used intermediates in the pharmaceutical chemistry to synthesize many biological active compounds. Ring fusion and addition reactions to their double bonds are a member of the most important reactions for Schiff’s bases. In this respect, it is intended in this work to benefit these opportunities in order to make compounds containing amino acids in their structure. Amino acids as natural products play a central role both as building blocks of proteins and as intermediates in metabolism. As main elements of proteins, the 20 essential amino acids expand an enormous diversity of chemical functionality [10]. Amino acid derivatives have been also indicated as essential component of drugs and exhibit antibacterial [11], antifungal [12], anticancer [13] and antibodies [14]. For example, the best known penicillin and cephalosporin comprise in their structures L-2-Aminoadipic acid, L-cysteine, and L-valine [15]. Therefore, the aimed compounds are expected to be potent antibacterial active compounds.

The aim of present work is to synthesize L-amino acid derivatives via Schiff’s bases which are derived from 2-aminopyridine, benzaldehyde or 4-chlorobenzaldehyde. Moreover, the antibacterial activity of the final compounds has to be evaluated against a Gram positive and a Gram negative bacteria (Staphylococcus aureus) and (Klebsiella pneumonia) respectively.

EXPERIMENTAL

Analytical instruments
Melting points were determined with Chachan, MLP-01 melting point apparatus. FT-IR spectra were recorded with FTIR-8300 Schimadzu, by KBr disk technique. ¹H-NMR and ¹³C-NMR spectra were recorded with Spectrometer 400 MHz, Avance III 400, Bruker, Germany in DMSO-d6 as a solvent.

Synthetic methods
Synthesis of N-(4-monosubstitutedbenzylidene) pyridin-2-amine 3(A,B)
(0.01 mol) of benzaldehyde 1A or 4-chlorobenzaldehyde 1B and (0.01 mol, 0.94 g) of 2-aminopyridine 2 were dissolved in 15 ml absolute ethanol containing a drop of glacial acetic acid and refluxed for 10 h. The reaction mixture was then allowed to cool to room temperature; the formed solid was filtered, washed with (2 %) HCl solution, then with water. The product was recrystallized from ethanol and dried [16].
Synthesis of N-benzyldiene pyridin-2-amine (3A)

Color=Yellow, yield=67%, m.p.=91°C. FT-IR (cm⁻¹): υ(C=O)=1678, υ(C=C) arom.=1597, υ(N=C–H) arom.=3147, υ(N=C–H) aliph.=2970 and 2950.  

Synthesis of N-(4-Chlorobenzylidene) pyridin-2-amine (3B)

Color=Yellow, yield=64%, m.p.=80°C. FT-IR (cm⁻¹): υ(C=C)=1678, υ(C=C) arom.=1585, υ(N=C–H) arom.=3147, υ(N=C–H) aliph.=2962 and 2931.  

Synthesis of N-[α-chloro (4-monosubstitutedbenzyl)]-N-pyrid-2-yl benzamide 5(A,B)

0.005 mol of 3(A, B) was dissolved in 10 ml dry benzene and placed in a three necked round bottom flask. 0.005 mol, 0.6 ml of benzyl chloride was dissolved in 6 ml dry benzene and poured in a dropping funnel. Benzyl chloride solution was added dropwise to the reaction mixture at 60°C with continuous stirring within 1 h. Then the solvent was removed and the solid residue was filtered, washed with (2%) solution of sodium carbonate, then with water, recrystallized from ethanol and dried [16].  

Synthesis of N-(α-chlorobenzyl) -N-pyrid-2-yl benzamide (5A)

Color=White, yield=63%, m.p.=101°C. FT-IR (cm⁻¹): υ(C=O) amide=1685, υ(C=C) arom.=1600, υ(N=C–H) arom.=3240, υ(N=C–H) aliph.=2927 and 2843, υ(N=C–H) =775.  

Synthesis of N-[α-chloro (4-Chlorobenzyl)]-N-pyrid-2-yl benzamide (5B)

Color=White, yield=74 %, m.p.=117°C. FT-IR (cm⁻¹): υ(C=O) amide=1685, υ(C=C) arom.=1593, υ(N=C–H) arom.=3070, υ(N=C–H) aliph.=2987 and 2885, υ(N=C–H) =799.  

Synthesis of (2S)-2-N[(benzoyl-pyridin-2-yl-amino)-(4-Chloro-phenyl-methyl)]-L-aminoc acid 6-10 (A,B)

0.005 mol of 5(A, B) and (0.005 mol) amino acid were dissolved in 10 ml (2:1) 1,4-dioxane-water and refluxed for 4 h. The resulting mixture was cooled and few drops of water were added, crystals were separated out, filtered and washed with water. The product was then recrystallized from (2:1) 1, 4-dioxane-water and dried [17].  

Synthesis of N-[(benzoyl-pyridin-2-yl-aminoyl)-phenyl-methyl]-Glucose (6A)

Color=White, yield=72%, m.p.=75°C. FT-IR (cm⁻¹): υ(C=O) overlaps between C=O amide and C=O acid, υ(OH)=3294, υ(NH)=3275, υ(C–C) arom.=1608, υ(C–H) arom.=3074, υ(N=C–H) arom.=3213, υ(N=C–H) aliph.=2927 and 2850. ¹H-NMR (ppm): δ=1.2 (1H, s, NH), δ=3.4 (2H, s, CH₂), δ=4.1 (1H, s, CH), δ=7.2-8.4 (14H, m, =CH of 3 aromatic rings), δ=11 (1H, s, OH). ¹³C-NMR (ppm): δ=60.1 (1C, CH₂), δ=72.3 (1C, CH₃), δ=114.7-152.1 (17C, 3 aromatic rings), δ=165.9 (1C, CO amide), δ=167.3 (1C, CO acid).  

Synthesis of (2S)-2-N[(benzoyl-pyridin-2-yl-aminoyl)-phenyl-methyl]-L-alanine (7A)

Color=White, yield=72%, m.p.=76°C. FT-IR (cm⁻¹): υ(C=O) overlaps between C=O amide and C=O acid, υ(OH)=3294, υ(NH)=3275, υ(C–C) arom.=1608, υ(C–H) arom.=3062, υ(N=C–H) arom.=3209, υ(N=C–H) aliph.=2927 and 2850. ¹H-NMR (ppm): δ=1.1 (1H, s, NH), δ=1.4 (3H, d, CH₃), δ=3.5 (1H, q, CH), δ=4.0 (1H, s, CH), δ=7.1-8.3 (14H, m, =CH of three aromatic rings), δ=10.7 (1H, s, OH). ¹³C-NMR (ppm): δ=60.1 (1C, CH₂), δ=72.4 (1C, CH₃), δ=114.6-152.1 (17C, 3 aromatic rings), δ=165.9 (1C, CO amide), δ=167.3 (1C, CO acid).  

Synthesis of (2S)-2-N[(benzoyl-pyridin-2-yl-aminoyl)-phenyl-methyl]-L-phenylalanine (7B)

Color=pale white, yield=72%, m.p.=73°C. FT-IR (cm⁻¹): υ(C=O)=1678 overlap between C=O amide and C=O acid, υ(OH)=3298, υ(NH)=3275, υ(C–C) arom.=1608, υ(C–H) arom.=3062, υ(N=C–H) arom.=3209, υ(N=C–H) aliph.=2931 and 2858. ¹H-NMR (ppm): δ=1.1 (1H, s, NH), δ=3.5 (1H, q, CH), δ=4.0 (1H, s, CH), δ=7.1-8.3 (13H, m, =CH of three aromatic rings), δ=10.7 (1H, s, OH). ¹³C-NMR (ppm): δ=36.3 (1C, CH₃), δ=54.3 (1C, CH), δ=114.7-166.2 (23C, 4 aromatic rings), δ=167.3 (1C, CO amide), δ=173.2 (1C, CO acid).  

Synthesis of (2S)-2-N[(benzoyl-pyridin-2-yl-aminoyl)-phenyl-methyl]-L-phenylalanin (8A)

Color=pale white, yield=68%, m.p.=71°C. FT-IR (cm⁻¹): υ(C=O)=1678 overlap between C=O amide and C=O acid, υ(OH)=3294, υ(NH)=3275, υ(C–C) arom.=1608, υ(C–H) arom.=3074, υ(N=C–H) arom.=3209, υ(N=C–H) aliph.=2927 and 2854. ¹H-NMR (ppm): δ=1.2 (1H, s, NH), δ=3.1 (2H, d, CH₃), δ=3.5 (1H, trp, CH), δ=4.0 (1H, s, CH), δ=7.1-8.3 (19H, m, CH of four aromatic rings), δ=10.7 (1H, s, OH). ¹³C-NMR (ppm): δ=36.3 (1C, CH₃), δ=54.3 (1C, CH), δ=114.7-166.2 (23C, 4 aromatic rings), δ=167.3 (1C, CO amide), δ=173.2 (1C, CO acid).  

Synthesis of (2S)-2-N[(benzoyl-pyridin-2-yl-aminoyl)-phenyl-methyl]-L-phenylalanin (8B)

Color=pale yellow, yield=71%, m.p.=74°C. FT-IR (cm⁻¹): υ(C=O)=1678 overlap between C=O amide and C=O acid, υ(OH)=3298, υ(NH)=3275, υ(C–C) arom.=1608, υ(C–H) arom.=3074, υ(N=C–H) arom.=3209, υ(N=C–H) aliph.=2931 and 2854. ¹H-NMR (ppm): δ=1.2 (1H, s, NH), δ=3.2 (2H, d, CH₂), δ=3.5 (1H, trp, CH), δ=4.0 (1H, s, CH), δ=7.1-8.3 (18H, m, =CH of three aromatic rings), δ=10.7 (1H, s, OH). ¹³C-NMR (ppm): δ=36.3 (1H, s, NH), δ=3.5 (2H, d, CH₂), δ=3.5 (1H, trp, CH), δ=4.0 (1H, s, CH), δ=7.0 (1H, d, CH), δ=7.1-8.3 (14H, m, =CH of three aromatic rings), δ=10.7 (2H, s, OH).
Synthesis of (2S)-2-N-[(benzoyl-pyridin-2-yl-amino)-(4-Chloro-phenyl)-methyl]-L-asparagine (9B)

Color=Pale white, yield=72 %, m.p.=77°C. FT-IR (cm⁻¹): υ(C=O)=1678 overlap between C=O amide and C=O acid, υ (OH)=3298, υ (NH)=3275, υ(N=C) arom.=1608, υ(C–H) arom.=3062, υ (N=–C–H) arom.=3209, υ (C–H) aliph.=2927 and 2854. ¹H NMR (ppm): δ=1.1 (1H, t, J=7 Hz, H3), δ=3.3 (3H, CH3), δ=7.0-8.2 (14H, m, CH, CH=C=O), δ=10.645 (1H, s, OH). ¹³C-NMR (ppm): δ=3.6 (1C, CH3), δ=54.3 (1C, CH2), δ=114.7-166.2 (17C, 3 aromatic rings), δ=167.3 (2C, C=O amide), δ=173.2 (1C, C=O acid).

Antibacterial activity

The antibacterial activity was performed according to the disc diffusion method [19]. Hereby the prepared agar and petri dishes were sterilized by autoclaving at 121°C for 15 min. The agar was inoculated uniformly from the broth culture of the tested microorganisms. In the solidified medium, suitably spaced apart holes were made 6 mm in diameter. These holes were filled with 5, 10, 25 and 50 μg/ml of the synthesized compounds dissolved in DMSO as a solvent. The plates were incubated at 37°C for 24 h.

RESULTS AND DISCUSSION

Chemistry

Scheme 1 shows the synthetic route starting from 2-aminopyridine 2 and benzaldehyde 1A or 4-chlorobenzaldehyde 1B towards aimed compounds 6-10 (A,B).

Schiff’s bases 3(A,B) were prepared through acid catalyzed condensation reaction of 2-aminopyridine 2 with benzaldehyde 1A or 4-chlorobenzaldehyde 1B in ethanol [16]. The formed Schiff’s bases may be a mixture of E/Z diastereomers. The condensation reaction follows an addition-elimination mechanism described below [18].

In the FT-IR spectrum the formation of 3(A,B) compounds are indicated by appearance of C=N absorption band at 1678 cm⁻¹, disappearance of carbonyl group absorption band at 1689 and 1699 cm⁻¹ and disappearance of the NH₂ absorption band at 3300 cm⁻¹ for symmetric and 3443 cm⁻¹ for asymmetric bands, respectively.

Schiff’s bases are nucleophiles and weak bases, so they do not react with simple allyl, alkyl or benzyl halides, but they react smoothly with the relatively more reactive acid halides such as benzoyl chloride [16]. The reaction of N-(4-monosubstituted benzylidene) pyridin-2-amine with benzoyl chloride yielded N-[chloro-(4-monosubstituted-phenyl)-methyl]-N-pyridin-2-yl-benzamide 5(A,B). The saturation of the double bond produced a new asymmetric carbon atom (stereogenic center) and therefore the product might be a racemic mixture of the two enantiomers. The reaction may follow a nucleophilic addition mechanism as follows [18].
In the FT-IR spectrum, the formation of compounds 5(A,B) are indicated by appearance of C–Cl absorption band 775 and 759 cm⁻¹, appearance of carbonyl group absorption band at 1685 cm⁻¹ and 1681 cm⁻¹, and disappearance of C=N absorption band at 1678 cm⁻¹.

(2S)-2-N-[(benzoyl-pyridin-2-yl-amino)-(4-monosubstituted-phenyl)-methyl]-L-amino acid 6-10 (A,B) have been synthesized by reaction of amino acid with N-[chloro-(4-monosubstituted-phenyl)-methyl]-N-pyridin-2-yl-benzamide in 2:1 of 1, 4-dioxane-water as a solvent [17]. The product now contains two chiral centers, the original carbon and the new generated carbon atom resulted from amino acids. That means the product is a mixture of diasteriomers. The reaction mechanism may follow a SN₂ mechanism described below [18,19].

FT-IR spectra of compounds 6-10 (A,B) show an appearance of a band at 1678 cm⁻¹, a band at 3275 cm⁻¹, and a band in the range of (3294-3302) cm⁻¹, which refer to the carbonyl amide, overlapped with the carbonyl carboxylic acid, NH-, and OH- groups, respectively. Also the disappearance of the absorption band at the range of (659-705) cm⁻¹ for chloride confirms the displacement of chloride with the amino acid.

In ¹H-NMR spectra of 6-10 (A,B) compounds the following proton assignments are noticed: a singlet signal at δ=0.9-1.2 ppm for NH proton, a singlet signal at δ=10.6-11.0 ppm for hydroxyl proton and multiplet signals at δ=7.0-8.2 ppm for benzene, benzoyl and pyridine protons.

In ¹³C-NMR spectra of 6-10 (A,B) compounds the following carbon assignments are detected: a signal at δ=114.7-166.2 ppm for benzene, benzoyl and pyridine aromatic rings, a signal at δ=165.9-167.3 ppm for carbonyl group amide and a signal at δ=167.2-173.2 ppm for acidic carbonyl group.

Antibacterial activities

The antibacterial activities were determined by measuring the diameter of the empty region around the wall (Inhibition zone). Results of preliminary screening tests are visualized in Figures 1 and 2. Generally, The synthesized compounds 6-10 (A,B) showed a lower antibacterial and moderate activity in comparison with Meropenem as a reference against both *Staphylococcus aureus* and *Klebsiella pneumonia*. From the obtained data in Figures 1 and 2, it is found that L-alanine compound 7B caused the highest activity against both types of the studied bacteria, but only when the concentration reaches 50 μg/ml. The glycine compound 6A caused a higher activity against *K. pneumonia* and moderate activity against *S. aureus*, when the concentration reaches 50 μg/ml. In principle, some compounds exhibited no activity, when the concentration was very low (5 and 10 μg/ml), as seen in compounds 6B, 7A, 7B, 8A, 9A, and 10A. Tested compounds could kill bacteria by destroying the cell wall. This can be explained by covalently binding to penicillin-binding proteins (PBPs) involved in the biosynthesis of mucopeptides in bacterial cell walls and this lead to rapid bacterial cell death. Bactericidal effects result through inhibition of cellular growth, division and the loss of cell wall integrity, which might cause cell wall lysis [20,21].

![Figure 1: Antibacterial effect of compounds 6-10 (A,B) on *Staphylococcus aureus*, compound concentration=5, 10, 25 and 50 μg/ml dissolved in DMSO at 37°C for 24 h, M=Meropenem as reference](image-url)
Glycine-6B and L-alanine-7B derivatives were the best candidates among the synthesized compounds, in which 4-chlorobenzaldehyde was used as starting material. Fundamentally, structural variation and substitution were important factors to be effective on weather Gram-positive or negative tested bacteria. Furthermore, it is clear, that more structural substitution and variation will be investigated and experimented to reach more favourable results.

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